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IN THE UNITED STATES PATENT AND TRADEMARK OPPICE

Applicants: Coppens et al.

Serial No.: 08/898,736

Filed:

July 23, 1997

Title:

PROCESS FOR THE PREPARATION OF MALTED

Group Art Voit: 1761

Braminer: C. Sherrer

CERTIFICATE OF FACSIMILE

I hereby certify that this paper for 08/898,736 is being facaimile transmitted to the U.S. Patent and Trademark Office at fax number 703 305 3602, on this dage.

Date Registration No. 30-192 Attorney for Applicant

SUPPLEMENTAL DECLARATION OF THE COPPERS TROSE 37 CPR 1.132

Honorable Commissioner of Patents and Trademerks Washington, D.C. 20231

Dene Bir:

- I, Theo Coppens, pursuant to 37 C.F.R. \$1.132, declare as Icilows:
- I am one of the inventors for the above-identified pacent application.
- In 1999, I asked Prof. C. Michiels, Professor of the Faculty of Agricultural and Applied Biological Sciences at Racholicke Universiteit Leuven in Belgium, to conduct the following experiments under my supervision to determine whether the medium and growth conditions described in Gyllang st al. would provide entivated spores. Those experiments and their results were first reported in my Declaration eigned on

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ALLY. DKt-NO. 61944

July 9, 1999. A more detailed explanation of those experiments and their results is presented herewith.

Materials and Methods

- 3. Chemicals and Media. Peptone, Yenst Extract and Rotato Dextrose Agar (PDA) were obtained from Unipath (Rampshire, United Kingdom). Dextrose was obtained from Merck-Belgolabo (Leuven, Belgium). Peptons, Yeast Extract and Dextrose madium was prepared according to Raiser at al. (1994). Peptons (2t w/v), Yeast Extract (1t w/v) ware dissolved in deionised water and sterilized at 121°C for 20 minutes. The pH of the obtained medium was 6.4.
- 4. Fungal Strains: Cultivation and Praparation of Culture Romanenate. The strains Rhisopus orysee ATCC 9363. Asporgillus fumigatus CBS 148.89 and Asporgillus amstelodami VTTD-76035 were obtained from respectively the American Type Culture Collection (ATCC, Manageas, VA, USA), Centralbureau voor Schimmelculturos (CBS, Banes, The Netherlands) and VTT (Technical Research Centre of Finland, Espoo, Finland) Culture collections. The strains were grown on PDA at 26°C. Seven days old sporulating cultures on PDA served as the starting material for culturing the fungi as described by Cyllang Gt al. (1977). For each strain a loopfull of material taken from the seven days old sporulating oulture on FDA was inoculated in a tissue culture flask containing 225 ml of Peptone, Yeast Extract and Dextrose medium. The culture was grown for 3 weeks at 20°C. After the oultivation period the entire culture was homogenized by vigorously shaking the content of the tissue oulture flask.

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bornessate. Activated spores were defined as described in the current patent application as 'being significantly more swollen than the dormant size, the size of the spores being increased by a factor preferably between 1.2 and 10 over the dormant spore size and/or having one or more germ tubes per spore. Three different samples of 0.2 ml of the culture homogenate were examined microscopically. Swolling of the spores was verified by measuring the spores at a magnification of 1250x by means of an eyepiede graticule micrometer. Activation was reported as a percentage of the spore population, determined by microscopic count. Therefore, the spore population was quantified by means of a Thoma counting chamber at magnification of 120x (Carl Zeiss, Jena, Germany). At least 100 spores per sample were evaluated.

Results

o. Analysis of spore activation. The dormant size of various fungal spores is described by Pict and Hocking (1957). According to this reference, the sporangiospores of Rhisopus orysas are of variable shape, ellipsoidal to broadly fusiform or irregularly angular, commonly 5.0 - 8.0 µm long, the condicepores of Aspergillus amstelodami-are apherical to subspheroidal with 4.0 - 5.0 µm dismeter; the condicepores of Aspergillus fumigatus are spherical to subspheroidal with 2.5 - 3.0 µm dismeter. Our own observations of dormant spores of the three tested strains were in agreement with the description given by Fitt and Bocking (1987). Accordingly, we dofined activated spores of Rhisopus crysae ATCC 9363, Aspergillus fumigatus CBS 148.89 and Aspergillus amstelodami VTT D-76035 as having respectively a size of more than 9.6 µm,

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6.0 µm and 3.6 µm and/or one or more germ tubes per spore.

Figure 1 shows some microphotographs of dormant, swellen and activated spores of Rhisopus oryses ATCC 9363.



Figure 1. Activated spores obtained by treatment as described in the current patent application (magnification 720%): A. dormant spores; S. swellen spores with one activated (AC) spore, i.a. Significantly more swellen than the dormant size; C. activated spores significantly more swellen than the dormant size; dormant size and having one or more germ tubes per spore.

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The results of the analysis of spore activation in the culture homogenates immediately after homogenisation (0 time) under the procedure of Gyllang et al. (1977) are presented in Table I.

Table I. Spore Activation at 6 Time in the Culture Econogenates.

	* ROSTER SETTIMATED
Rhisopus oryses ATCC 9363	¢
Aspergilius fumigatus CBS 148.89	Q
Aspergillus amacelodami VTT D-76015	0

Further activation of spores in the culture homogeness was analyzed after 6 hours incubation of the culture homogeneous at 20°C or 42°C, although this deviates from the procedure of Gyllang et al. (1977). In this procedure no incubation period is preseribed between preparation of the homogenete and incoulation of the barley. The results are presented in Table II.

Table II. Spore Activation After 6 Hours Insubstica in the Culture Homogenates.

	* spores	activated
	Indubation	Incubation
Rhizopus orysee ATCC 9363	0	0
Aspergillus fumigatus CBS 148.88	•	0
Aspergillus amsteledami VTT D-76035	3	0

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In contrast, treatment of Rhizopus oryses ATCC 9367 spores as described in the current patent application resulted in a high level of activation of the spore as more than 90% of the spores had a size of more than 9.6 μm and/or had one or more gottm tubes per spore.

- 7. Conclusions. Culture homogenates of Rhisopus orysee ATCC 9363, Aspergilius fumigatus CES 146.69 and Aspergilius emstelodami VTT D-76036 prepared according to Gyllang et al. (1977) do not contain activated spores. This experiment shows that successful activation depends on incubation of dormant spores for a sufficient time at a suitable temperature and in a suitable medium. In the spore suspension as prepared by Gyllang et al. (1977) the medium is an exhausted growth medium that does not provide the suitable conditions for spore activation, and the spores are not incubated for a sufficient time at a suitable temperature.
- 8. <u>Abbreviation used</u>. FDR, Forato Dextrose Agar; ATCC; American Type Culture Collection; CBF, Centrasibureau voor Schimmeliultures; VTT, Technical Research Centre of Pinland; Ac, sotivated.

9. Reforences.

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The undersigned, being warned that willful felse statements and the like are punishable by fine or imprisonment, or both (18 U.S.C. E1091) and may jeopardize the velidity of the application or any patent issuing thereon, bereby declares that the above statements made of my own knowledge are true and that all statements made on information and belief are believed to be true.

Theo Coppens

Date: 03/03/2000

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